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#### (57) Abstract

The present invention is based upon the finding that porcine endogenous retroviruses exist in two different subtypes, which we have termed PERV-A and PERV-B. The differences are reflected in sequence divergence in the envelope genes, and these differences may be used to provide nucleic acid and antibody probes which can distinguish between the two subtypes. This allows patterns of subtype transmission between cells, particularly porcine to human cells, to be monitored.

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# DETECTION OF RETROVIRAL SUBTYPES BASED UPON ENVELOPE SPECIFIC SEQUENCES

The present invention relates to methods and products for the detection of porcine endogenous retroviruses.

There is currently much interest in the development of xenotransplantation of organs to meet the shortage of human organs available for transplant. Considerable progress has been made in developing transgenic animals, particularly pigs, whose organs have been modified to remove immunogenic surface antigens and/or to present human antigen, or to inhibit components of the human immune system. However while progress has been made on the immunological problems of xenotransplantation, relatively little research has been conducted on the risk of infection being transmitted to an organ recipient by the presence of endogenous pathogens in the donor organ.

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- 15 Recently, Patience et al, Nature Medicine, 1997, 3;282-286, reported the results of a study of pig endogenous retroviruses (PERVs) in porcine cell lines. The authors demonstrated that two different pig kidney cell lines, PK15 and MPK, produced endogenous retroviruses and the PK15 retroviruses were capable 20 of infecting a human cell line (kidney 293 cells). of the protease and reverse transcriptase genes of the retroviruses infecting these cell lines showed that there was about 95% sequence similarity at the amino acid level between isolates from the two cell lines. This information was used 25 to design nucleic acid primers for the analysis of DNA from porcine tissue and the authors demonstrated that multiple PERV related sequences existed in such tissue and were expressed. The primers were specific for porcine PERVs and did not detect sequences in human or murine cells.
- WO97/21836, published on 19 June 1997, describes three porcine retrovirus isolates. These isolates are currently described as PERV-A and PERV-C, with SEQ ID NO:1 and SEQ ID NO:3 of

WO97/21836 being of the PERV-C type, and SEQ ID NO:2 being of the PERV-A type.

WO97/40167, published on 30 October 1997, describes a retrovirus isolate from the PK-15 porcine cell line. This isolate is currently described in the art as being of a PERV-B type. Figure 3 of WO97/40167 sets out a sequence with 3 open reading frames indicated to be the gag, pol and env genes of the retrovirus. Figure 1 of WO97/40167 sets out a shorter sequence with a 3' end which extends into the 5' region of the env gene. There are differences between the 3' end of Figure 1 and the corresponding region of Figure 3. The differences are attributed in WO97/40167 to improvements in carrying out and analysing the sequence obtained.

#### Disclosure of the invention.

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15 Prior to the present invention, it had not been appreciated that PERVs existed in different subtypes. Prior to the publication of WO97/21836 and WO97/40167 we surprisingly identified two subtypes of this virus, which we designated PERV-A and PERV-B. More surprisingly, although the majority of individual isolates from the PK15 cell line are PERV-A isolates (29/32 tested), our initial data indicated that human 293 cells infected with the virus are exclusively or almost exclusively of the PERV-B subtype. Thus although the primers used by Patience et al are capable of detecting numerous PERV sequences in porcine tissue and cell lines, these primers do not distinguish between the two subtypes of PERV.

In the light of the present invention we believe that the sequence of Figure 1 of WO97/40167 is derived from a PERV-A isolate, since the Figure 1 sequence in the region of difference is substantially similar to the corresponding portion of the PERV-A isolate described herein.

In a first aspect the present invention thus provides an

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isolated nucleic acid probe, said probe being capable of hybridising to the PERV-B env gene under conditions in which said probe is substantially unable to hybridise to the PERV-A env gene. This is referred to below as a PERV-B specific probe (or "primer" or "oligonucleotide"). The terms "probe", "primer" and "oligonucleotide" are used synonymously.

In a second aspect, the invention provides an isolated nucleic acid probe, said probe being capable of hybridising to the PERV-A env gene under conditions in which said probe is substantially unable to hybridise to the PERV-B env gene. This is referred to below as a PERV-A specific probe (or "primer" or "oligonucleotide").

Although the *env* gene sequences are shown as the positive strand, it is to be understood that probes of the invention may be directed to either strand where integrated or cDNA retroviral sequences are to be detected. Where retroviral RNA is to be detected, a probe capable of hybridising to the positive strand is required (in the case of PCR initially to make cDNA).

In a further aspect, the invention provides a pair of primers suitable for conducting a polymerase chain reaction, at least one of said primers being a nucleic acid as defined above specific for the PERV-A or PERV-B genes. The probes and primers of the invention may be used in a method of detecting retroviruses in a sample of porcine or human tissue. Such tissue includes primary porcine tissue and human cell lines which have been cultivated in the presence of a porcine cell line, or human tissues which are from a human patient who has received a xenotransplant. Nucleic acid (e.g. mRNA, total RNA, DNA or total nucleic acid) from the tissues or cells may be probed directly or if desired retroviral sequences may be amplified using primers suitable for amplifying retroviral sequences in general (e.g. LTR primers) prior to detecting PERV env sequences of the invention, thus allowing those of

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skill in the art to distinguish between the PERV-A and PERV-B subtypes. The nucleic acid may be present in a sample comprising human or porcine tissue or cells, or may be cloned nucleic acid from such sources.

5 The differences between the two genes is reflected by changes to the env proteins, and these differences are believed to include differences to antigenic determinants (referred to herein as epitopes) in the two subtypes of proteins, which thus allows the development of antibodies which are capable of binding to an epitope on the PERV-B env protein under conditions where they are substantially unable to bind to the PERV-A env protein, and vice versa. These antibodies may be used in a method of detecting the presence of a pig endogenous retrovirus in porcine or human tissue or cell lines, thus allowing those of skill in the art to distinguish between the PERV-A and PERV-B subtypes.

# Detailed Description of the Invention.

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Our prototype isolate of the PERV-A env gene region is shown in SEQ ID NO. 1, and the envelope polypeptide encoded by nucleotides 211 to 2190 of SEQ ID NO. 1 is shown as SEQ ID NO. 2. For the purposes of the present invention, the PERV-A env gene is at least 80%, preferably at least 90% and more preferably at least 95% homologous to the coding sequence of SEQ ID NO. 1. Homologous sequences include those which encode the same polypeptide shown in SEQ ID NO:2 but differ from SEQ ID NO:1 due to the degeneracy of the genetic code.

The percentage homology (also referred to as identity) of DNA sequences can be calculated using commercially available algorithms, such as Lasergene software from DNASTAR Inc or the algorithm GAP (Genetics Computer Group, Madison, WI). GAP uses the Needleman and Wunsch algorithm to align two complete sequences that maximizes the number of matches and minimizes the number of gaps. Generally, the default parameters are

used, with a gap creation penalty = 12 and gap extension penalty = 4. Use of either of the terms "homology" and "homologous" herein does not imply any necessary evolutionary relationship between compared sequences, in keeping for example with standard use of terms such as "homologous recombination" which merely requires that two nucleotide sequences are sufficiently similar to recombine under the appropriate conditions.

Similarly, our prototype isolate of the PERV-B env gene region is shown in SEQ ID NO. 3, and the envelope polypeptide encoded by nucleotides 911 to 2881 of SEQ ID NO. 3 is shown as SEQ ID NO. 4. For the purposes of the present invention, the PERV-B env gene is at least 80%, preferably at least 90% and more preferably at least 95% homologous to the coding sequence of SEQ ID NO. 3. Homologous sequences include those which encode the same polypeptide shown in SEQ ID NO:4 but differ from SEQ ID NO:3 due to the degeneracy of the genetic code.

An alignment of SEQ ID NO. 1 and SEQ ID NO. 3 is shown as Figure 1.

- The PERV-B specific probe of the invention is preferably derived from the 5' end of the env gene of PERV-B, particularly from the region of PERV-B corresponding to nucleotides 1000 to 2500 of the SEQ ID NO. 3 isolate. More preferably the region corresponds to nucleotides 1100 to 1900.
- It is to be understood that "derived" means conceptually derived, and physical isolation of the nucleic acid from the gene (as opposed to, for example, de novo synthesis) is not necessary.
- Specific PERV-B probes include oligonucleotides consisting of a contiguous sequence of from 10 to 40 nucleotides of a PERV-B isolate derived from the sequence of SEQ ID NO:3 from 1000 to 2500, preferably 1100 to 1900, or the complement thereof.

Such oligonucleotides include SEQ ID NO:7 (1376-1395 of SEQ ID NO:3) and SEQ ID NO:8 (complement of 1620-1639 of SEQ ID NO:3) shown in Example 3 below comprise 8 and 14 differences respectively in their sequences and the corresponding regions of SEO ID NO:1 as follows:

PERV-B 5' TTCTCCTTTGTCAA--TTCCGG 3' (SEQ ID NO:7)

\* \* \*

PERV-A 5' TACTCTTTTGTTAACAATCCTA 3' (SEQ ID NO:9)

and:

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PERV-A 5' TATTCTGAGGCGCGAATAGT 3' (SEQ ID NO:10)

Similarly, the PERV-A specific probe of the invention may be derived from the regions shown in Figure 1 which correspond to the abovementioned preferred and most preferred regions of PERV-A. Thus PERV-A specific probes include oligonucleotides consisting of a contiguous sequence of from 10 to 40 nucleotides of a PERV-A isolate derived from the sequence of SEQ ID NO:1 from 300 to 1809, preferably 400 to 1209, or the complement thereof.

Thus for example such oligonucleotides include SEQ ID NO:5 (742-760 of SEQ ID NO:1) and SEQ ID NO:6 (complement of 1082-1101 of SEQ ID NO:1) shown in Example 3 below. These comprise 10 and 21 differences respectively in their sequences and the corresponding regions of SEQ ID NO:3.

By "differences", it is meant substitutions, deletions and insertions. As can be seen from Figure 1, the primers of SEQ ID NOs:5-8 include between them all these differences from the corresponding portions of the reference isolate.

30 The above-mentioned probes may additionally include, at their

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3' and/or 5' termini, linker sequences (typically of from 3 to 8 nucleotides) of non-PERV-B or -A sequence. Linker sequences include those containing a restriction enzyme recognition sequence allowing the oligonucleotides to be introduced into or excised from a cloning or expression vector.

Nucleic acid probes of the invention may be obtained by first of all comparing the PERV-A and PERV-B sequences of Figure 1 (or of other PERV-A and PERV-B isolates) and regions of the sequences which are sufficiently different to provide specific probes determined. This may be done by any suitable means, for example by calculating the predicted Tm of a probe when annealed to a specific region of the PERV-A or PERV-B sequences using a suitable algorithm or empirically by experiment. When by experiment this can be achieved by blotting the PERV-A and PERV-B sequences onto a nitrocellulose filter and probing the filter with a labelled putative probe under hybridising conditions. Probes of the invention will be able to hybridise to the PERV sequence of choice and not to the other PERV sequence under those conditions. Thus a PERV-B specific probe of the invention will be capable of hybridising to the sequence of SEQ ID NO:3 under conditions in which the probe does not hybridise to SEQ ID NO:1. Similarly, a PERV-A specific probe of the invention will be capable of hybridising to the sequence of SEQ ID NO:1 under conditions in which it does not hybridise to SEQ ID NO:3.

Hybridisation conditions will be selected to be commensurate with the size of the probe and can be determined by reference to standard text books such as Sambrook et al, Molecular Cloning, 1989, Cold Spring Harbour.

It will be understood by those of skill in the art that hybridisation conditions will vary depending upon whether a probe of the invention is hybridised to nucleic acid fixed to a solid support or is hybridised to a target nucleic acid in a liquid phase. In the case of the former (eg Southern or

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Northern blotting) a probe of the invention will be annealed under low stringency conditions and subsequently washed under high stringency conditions such that the probe will remain annealed to its target PERV sequence and not to the corresponding sequence of the other subtype. Where a probe of the invention is for use as a PCR primer annealing conditions will be selected in accordance with standard protocols such that the probe will hybridise to its target subtype nucleic acid and not to non-target subtype nucleic acid. Thus it will be understood that reference to hybridisation of a probe to target nucleic acid includes hybridisation achieved by blotting and washing on a solid phase as well as annealing in a liquid phase. In either case, the person of skill in the art will be able to test using routine skill and knowledge whether any selected sequence derived from a PERV-B env gene is able to hybridise to the PERV-B env nucleic acid under conditions in which it is substantially unable to hybridise to PERV-A env nucleic acid, and vice versa.

One way to calculate Tm of a probe is by reference to the formula for calculating the Tm of probes to a homologous target sequence. This formula is Tm(°C) = 2(A+T) + 4(G+C) -This will provide the Tm under conditions of 3xSSC and 0.1% SDS (where SSC is 0.15M NaCl, 0.015M sodium citrate. pH 7). This formula is generally suitable for probes of up to 30 nucleotides in length. In the present invention, this formula may be used as an algorithm to calculate a nominal Tm of a probe for a specified sequence based upon the number of matches to its PERV target (e.g. PERV-B) sequence and PERV non-target sequence (e.g. PERV-A). For example, for the probe of SEQ ID NO:7 has a Tm of ((2x11) + (4x9) - 5) = 53°C. sequence of SEQ ID NO:7 is derived from SEQ ID NO:3 and thus will have this Tm when used as a probe for this sequence, subject to the usual experimental error. However when SEQ ID NO:7 is used as a probe for the corresponding region of SEQ ID NO:1 (represented above as SEQ ID NO:9), the calculated Tm will be ((2x9) + (4x5) - 5) = 33°C, based on counting the

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number of matches. (Since for the purposes of the present invention the above formula is used as an algorithm, the actual Tm of probes when hybridised to non-complementary targets which do not exactly match the probe sequence may or may not correspond to the calculated value.)

Thus in a preferred aspect, a PERV-B specific probe will have a Tm (calculated as above) for SEQ ID NO:3 which is at least 5°C higher than for SEQ ID NO:1, and vice versa for a PERV-A specific probe. Preferably the difference is at least 8°C, more preferably at least 10°C, at least 15°C or at least 20°C.

The above formula generally useful for probes up to 30 nucleotides in length, but since it is used simply as an algorithm in the present invention, it may be extended to longer probes, for example up to 40 or even up to 50 nucleotides in length.

Suitable conditions for a probe to hybridise to a PERV target sequence may also be measured experimentally. Suitable experimental conditions comprise hybridising a candidate probe to both SEQ ID NO:1 and SEQ ID NO:3 on a solid support under low stringency hybridising conditions (e.g. 6xSSC at 55°C), washing at reduced SSC and/or higher temperature, for example at 0.2xSSC at 45°C, and increasing the hybridisation temperature incrementally to determine hybridisation conditions which allow the probe to hybridise to SEQ ID NO:1 but not SEQ ID NO:3, or vice versa, as the case may be.

Although the hybridisation conditions used to distinguish between the PERV-B and PERV-A env genes should also be sufficient to distinguish over other "background" sequences present in human or porcine cells (particularly human and porcine genomic and mitochondrial sequences), it is also desirable that the probes do not, under such conditions, hybridise to such background sequences. This may also be determined by experiment, for example by blotting the probes

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to a solid support which carries at separate loci SEQ ID NO:1, SEQ ID NO:3 (for example cloned in plasmids), human total DNA and porcine total DNA.

The size of the probe may be selected by those of skill in the art taking account of the particular purposes the probes are to be used. Probes may be for example from 10 to 1000 nucleotides (or base pairs), e.g. from 50 to 500, such as from 200 to 500 nucleotides or base pairs. This size range is particularly suitable for Southern blots. However for some purposes, for example PCR, short oligonucleotide probes are preferred, generally in the size range of from 10 to 40 nucleotides in length, preferably 12 to 25 and more preferably from 18 to 24 such as 20, 21 or 22 nucleotides.

The probes may be labelled with a detectable label, including a radionuclide such as  $^{32}P$  or  $^{35}S$  which can be added to the probe using methods known per se in the art. The probe may alternatively carry a non-radioactive label such as biotin.

Generally, probes will be prepared by stepwise chemical synthesis, which is widely available commercially.

Recombinant production of probes is also possible. Probes may be DNA or RNA, and may contain or consist of synthetic or modified nucleotides. A number of different types of modification to oligonucleotides are known in the art. These include methylphosphonate and phosphorothionate backbones, addition of acridine or polylysine chains at the 3' and/or 5' ends of the molecule. For the purposes of the present invention, it is to be understood that the probes and primers described herein may be modified by any method available in the art.

A preferred method of detection is by the polymerase chain reaction (PCR). This will involve PERV-B or PERV-A primer pairs, at least one of which is directed to PERV-B or PERV-A env gene sequences, the polarity of the probes being such that

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the region between them is amplified when the PCR is performed. At least one of each pair of PERV-A and/or PERV-B primers will be specific for its target PERV sequence. The other member of each pair may be targeted to non-env sequence or env sequence common to PERV-A and PERV-B. Preferably both members of a primer pair are specific for their target PERV sequence. Desirably the probes will be selected to amplify a region of the PERV-A and PERV-B of a convenient size to detect, such as between about 50 and 500, preferably between 150 and 400 nucleotides.

Where pairs of PERV-A and PERV-B primers are used in conjunction with each other, it is preferred that the primer pairs are selected such that different size PERV-A and PERV-B products are produced. Preferably the difference in size is at least from 5 to 50 base pairs, such as from 10 to 25 base pairs, so that detection of the products by electrophoresis on agarose gels by ethidium bromide staining may be conveniently carried out.

The methods of the invention which allow the PERV-A and PERV-B subtypes to be distinguished are useful in following the transmission of these viruses from porcine cells to other cell types, particularly human cells. In addition, the probes may be used to clone and characterize the different endogenous proviruses of pigs. Specific proviruses can be characterised by both their sequences and the genomic flanking sequences, and thus a map of the chromosomal locations of the viruses may be determined. The ability to distinguish between PERV-A and PERV-B proviruses will facilitate studies of the porcine endogenous retroviruses which might pose a threat to humans in a transplant setting.

The PERV-A and PERV-B nucleic acid sequences of the invention are novel and thus in a further aspect of the invention there is provided an isolated nucleic acid consisting essentially of the PERV-A or PERV-B env gene coding sequence, or a fragment

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thereof which is capable of hybridising to the PERV-B env gene under conditions in which said probe is substantially unable to hybridise to the PERV-A env gene, or vice versa. Vectors which comprise such sequences form a further aspect of the invention. The vector may be for replication of the sequence or for expression of the sequence in a suitable host cell. In such a case the vector will comprise a promoter operably linked to the env sequence, the promoter being compatible with the host cell which may be, for example, bacterial, e.g. E.coli, yeast, insect or mammalian, e.g. a CHO cell or a human cell line.

The env gene may be expressed in such a cell and recovered from the cell in substantially isolated form.

The differences in the PERV subtypes also allow the production of antibodies which can distinguish between the two subtypes. In a manner analogous to the production of probes, the sequence differences between the proteins of SEQ ID NO. 2 and SEQ ID NO. 4 can be examined, and suitable epitopes which reflect these differences determined using computer algorithms or by epitope scanning techniques. Monoclonal antibodies raised against these epitopes may be used to detect the presence of the PERV-A and/or PERV-B subtypes in a specific manner.

In a manner analogous to the nucleic acid probes, the antibodies are preferably directed to epitopes in the N-terminal region of the PERV-A and PERV-B env proteins, particularly epitopes encoded within the preferred regions identified above.

For the purposes of the present invention the term antibody describes an immunoglobulin whether natural or partly or wholly synthetically produced. The term also covers any polypeptide or protein having a binding domain which is, or is homologous to, an antibody binding domain. These can be

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derived from natural sources, or they may be partly or wholly synthetically produced. Examples of antibodies are the immunoglobulin isotypes and their isotypic subclasses; fragments which comprise an antigen binding domain such as Fab, scFv, Fv, dAb, Fd; and diabodies.

It is possible to take monoclonal and other antibodies and use techniques of recombinant DNA technology to produce other antibodies or chimeric molecules which retain the specificity of the original antibody. Such techniques may involve introducing DNA encoding the immunoglobulin variable region, or the complementarity determining regions (CDRs), of an antibody to the constant regions, or constant regions plus framework regions, of a different immunoglobulin. See, for instance, EP-A-184187, GB 2188638A or EP-A-239400. A hybridoma or other cell producing an antibody may be subject to genetic mutation or other changes, which may or may not alter the binding specificity of antibodies produced.

As antibodies can be modified in a number of ways, the term "antibody" should be construed as covering any specific binding member or substance having a binding domain with the required specificity. Thus, this term covers antibody fragments, derivatives, functional equivalents and homologues of antibodies, including any polypeptide comprising an immunoglobulin binding domain, whether natural or wholly or partially synthetic. Chimeric molecules comprising an immunoglobulin binding domain, or equivalent, fused to another polypeptide are therefore included. Cloning and expression of chimeric antibodies are described in EP-A-0120694 and EP-A-0125023.

It has been shown that fragments of a whole antibody can perform the function of binding antigens. Examples of binding fragments are (i) the Fab fragment consisting of VL, VH, CL and CH1 domains; (ii) the Fd fragment consisting of the VH and CH1 domains; (iii) the Fv fragment consisting of the VL and VH

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domains of a single antibody; (iv) the dAb fragment (Ward, E.S. et al., Nature 341, 544-546 (1989)) which consists of a VH domain; (v) isolated CDR regions; (vi) F(ab')2 fragments, a bivalent fragment comprising two linked Fab fragments (vii) single chain Fv molecules (scFv), wherein a VH domain and a VL domain are linked by a peptide linker which allows the two domains to associate to form an antigen binding site (Bird et al, Science, 242, 423-426, 1988; Huston et al, PNAS USA, 85, 5879-5883, 1988); (viii) bispecific single chain Fv dimers (PCT/US92/09965) and (ix) "diabodies", multivalent or multispecific fragments constructed by gene fusion (WO94/13804; P. Holliger et al Proc. Natl. Acad. Sci. USA 90 6444-6448, 1993).

The reactivities of antibodies to an epitope in a sample may be determined by any appropriate means. Tagging with individual reporter molecules is one possibility. The reporter molecules may directly or indirectly generate detectable, and preferably measurable, signals. The linkage of reporter molecules may be directly or indirectly, covalently, eg via a peptide bond or non-covalently. Linkage via a peptide bond may be as a result of recombinant expression of a gene fusion encoding antibody and reporter molecule.

One favoured mode is by covalent linkage of each antibody with an individual fluorochrome, phosphor or laser dye with spectrally isolated absorption or emission characteristics. Suitable fluorochromes include fluorescein, rhodamine, phycoerythrin and Texas Red. Suitable chromogenic dyes include diaminobenzidine. Other reporters include macromolecular colloidal particles or particulate material such as latex beads that are coloured, magnetic or paramagnetic, and biologically or chemically active agents that can directly or indirectly cause detectable signals to be visually observed, electronically detected or otherwise recorded. These molecules may be enzymes which catalyse

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reactions that develop or change colours or cause changes in electrical properties, for example. They may be molecularly excitable, such that electronic transitions between energy states result in characteristic spectral absorptions or emissions. They may include chemical entities used in conjunction with biosensors. Biotin/avidin or biotin/streptavidin and alkaline phosphatase detection systems may be employed.

A radionuclide such as <sup>125</sup>I, <sup>111</sup>In or <sup>99m</sup>Tc may be attached to an antibody and these nuclides are useful in imaging target antigens in the body. Antibodies labelled with these labels may be used to examine xenotransplanted organs in a human recipient for the presence of PERVs as part of ongoing monitoring following transplantation.

Antibodies of the invention may be produced by conventional hybridoma technology, e.g by linking a peptide comprising a suitable epitope to a carrier protein, injecting the linked peptide into an animal such as a rat or rabbit, recovering the spleen and producing hybridoma cell lines which are screened against the peptide for specific binding. Antibodies may also be prepared by screening against synthetic libraries such as phage display libraries. Antibodies may also be made against the entire env protein or substantial parts thereof, and then screened individually against PERV-A and PERV-B env protein for specific binding to one or the other.

In one aspect of the invention a specific PERV-A antibody and a specific PERV-B antibody are used on parallel samples (or on the same sample where the two antibodies are labelled with different and distinguishable labels) to detect the presence of the two subtypes of retroviruses.

Antibodies specific for a PERV-B epitope will have at least a 100 fold higher affinity for that epitope than for the corresponding region (as indicated by alignments to the PERV-A

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sequence such as that of Figure 1) of the PERV-A env protein., and vice versa. Desirably both types of specific antibodies will not cross react to other proteins normally present in human and porcine cells (i.e. have at least a 100 fold higher affinity to its target epitope than to such other proteins).

The probes, primers and antibodies of the invention may be used in all aspects of the development of porcine organ (e.g. kidney, liver, heart, pancreas, including tissues and cells therefrom, such as pancreatic islet cells) xenotransplantation. Thus the probes, primers and antibodies may be used to monitor the inheritance of human tropic viruses, thus facilitating the breeding of pigs lacking these viruses, particularly the PERV-B subtype. The invention will also be useful in monitoring the expression of the viruses in pigs and humans.

The following examples illustrate the invention.

#### Example 1: Cloning of PERV-A and PERV-B Env sequences.

cDNA clones were obtained using the 3' RACE technique (Frohman and Martin Technique 1:165-170, 1989). Total RNA from PK15, MPK and 293 cells was reverse transcribed to produce cDNA using an adapter primer dT-Ri-Ro.

A fraction of cDNA from PK15 and MPK cells was amplified by the polymerase chain reaction (PCR) using the primer PL146 (5'ATCCGTCGGCATGCATAATACGACTCAC, SEQ ID NO:11) in combination with PL135 (5'CGATTCAGTGCTGCTACAAC, SEQ ID NO:12) or PL137 (5'CCCTTATAACCTCTTGAGCG, SEQ ID NO:13). Products of approximately 6.5 kb were digested with XhoI and SphI and cloned into SalI//SphI digested pGem3Zf(+). Positive clones were identified and sequenced.

30 A portion of cDNA from 293 cells was amplified by PCR using primer PL137 in combination with primer Ro. Products of

approximately 6.5 kb were isolated and digested with PstI and ligated with the pGem3Zf(+) plasmid digested with PstI and SmaI. After transformation into E.coli, positive clones were identified and sequenced.

Further clones were generated and sequenced from MPK and PK15 cDNA by amplification with primer PL147 (5'GTAATGCATGCTTCTATGGTGCCAGTCG, SEQ ID NO:14) in combination with either PL135, PL137 or PL148 (5'CTCTACGCATGCGTGTACGACTGTG, SEQ ID NO:15) and digestion of products with XhoI/SphI or SphI and cloning into appropriately digested pGEM3Zf(+).

Further clones were generated and sequenced from 293 cDNA by PCR amplification with primer PL147 in combination with either PL135, PL137 or PL149 (5'GTAATCGGGTCAGACAATGG, SEQ ID NO:16) and digestion of products with *EcoRI/PstI*, *PstI*, or *BamHI/EcoRI* and cloning into appropriately digested pGem3Zf(+).

Oligos dT-Ri-Ro and Ro come from Frohman and Martin (Technique 1:165-170,1989), PL146 is a modified version of Ro containing an additional SphI site, PL135 and PL137 were designed from the published PERV pol sequence (Tristan et al J.Virol 70:8241-8246, 1996 Genbank ID X99933), PL147 and PL148 are PERV LTR primers derived from the sequences of our initial 293 clones.

Analysis of the clones identified two distinct subtypes, which we have termed PERV-A and PERV-B. An alignment of the two subtype envelope gene sequences is shown in Figure 1.

Example 2: Frequency of full length PERV-A and PERV-B env gene isolation.

The frequency of the subtypes in pig and human cells was analysed and the results are as follows:

- 1. From pig PK-15 cells
  29/32 PERV-A 3/32 PERV-B
- 2. From human 293 cells infected with PK15 virus 0/18 PERV-A 18/18 PERV-B

#### 5 Example 3: Preparation of specific probes

#### 1. PCR

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Differences between the PERV-A and PERV-B subgroups allow the design of specific primers

PL170 TGGAAAGATTGGCAACAGCG (SEQ ID NO:5)

10 PL171 AGTGATGTTAGGCTCAGTGG (SEQ ID NO:6)

PL172 TTCTCCTTTGTCAATTCCGG (SEQ ID NO:7)

PL173 TACTTTATCGGGTCCCACTG (SEQ ID NO:8)

PL170+PL171 are predicted to give a 361 base pair band with PERV-A;

- PL172+PL173 are predicted to give a 264 base pair band with PERV-B. PCR studies with cloned plasmid DNA confirmed these prediction and showed no cross-amplification between the two primer pairs. Sequencing the respective RT-PCR products from RNA containing both viral RNAs shows amplification only of the sequences predicted from each primer pair.
  - 2. Southern blot probes.

The amplification products of PL170+PL171 (361 bp, PERV-A probe) and PL172+PL173 (264 bp, PERV-B probe) show no cross hybridisation on plasmid blots. Both have been used on genomic southern blots.

## Example 4: Host range studies

The host range specified by the cloned PERV *env* genes were examined using a Moloney murine leukemia virus (Mo-MLV) based vector to deliver the  $\beta$ -galactosidase (lacZ) indicator gene to different cell types (Tailor *et al* J.Virol. 67:6737-6741,

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1993). The TELCeB6 cell line (Cosset et al J.Virol. 69: 7430-7436, 1995) is derived from TE671 cells by stable transfection with CeB to supply the Mo-MLV gag-pol genes and carrying a modified lacZ gene (Ferry et al PNAS 88: 8377-8381, 1991) in proviral context introduced by infection using an amphotropic viral vector. The PERV env genes were introduced by transfection of TELCeB6 cells with expression constructs derived from pFBMOSALF (Cosset et al J.Virol, 69: 6314-6322, 1995) in which the PERV sequences, on XbaI-ClaI fragments, replace the corresponding Mo-MLV envelope sequence. Virus produced by transiently and stably transfected TELCeB6 cells were assayed for transfer of LacZ on 293, TE671 (human) and PK-15, PAE, ST-IOWA (pig) cells. Transfer of retroviral particles comprising the PERV-B envelope to human cells was demonstrated.

The infectious titre (LacZ positives/ml supernatant) was as follows:

	Pig	Mink	Human	Human
Virus	(ST-IOWA)	(Mu-1-lv)	(293)	(TE671)
PERV-A	2000	1000	300	2000
PERV-B	800	4000	800	700

# SEQUENCE LISTING

SEQ ID NO.	1: PERV-A.s	eq			
TCGAGTGGGT	GAGGCAGCGA	GCGTGGAAGC	AGCTCCGGGA	GGCCTACTCA	
GGAGGAGACT	TGCAAGTTCC	ACATCGCTTC	CAAGTTGGAG	ATTCAGTCTA	100
TGTTAGACGC	CACCGTGCAG	GAAACCTCGA	GACTCGGTGG	AAGGGACCTT	
ATCTCGTACT	TTTGACCACA	CCAACGGCTG	TGAAAGTCGA	AGGAATCCCC	200
ACCTGGATCC	ATGCATCCCA	CGTTAAGCCG	GCGCCACCTC	CCGATTCGGG	
GTGGAAAGCC	GAAAAGACTG	AAAATCCCCT	TAAGCTTCGC	CTCCATCGCG	300
TGGTTCCTTA	CTCTGTCAAT	AACTCCTCAA	GTTAATGGTA	AACGCCTTGT	
GGACAGCCCG	AACTCCCATA	AACCCTTATC	TCTCACCTGG	TTACTTACTG	400
ACTCCGGTAC	AGGTATTAAT	ATTAACAGCA	CTCAAGGGGA	GGCTCCCTTG	
GGGACCTGGT	GGCCTGAATT	ATATGTCTGC	CTTCGATCAG	TAATCCCTGG	500
TCTCAATGAC	CAGGCCACAC	CCCCGATGT	ACTCCGTGCT	TACGGGTTTT	
ACGTTTGCCC	AGGACCCCCA	AATAATGAAG	AATATTGTGG	AAATCCTCAG	600
GATTTCTTTT	GCAAGCAATG	GAGCTGCATA	ACTTCTAATG	ATGGGAATTG	
GAAATGGCCA	${\tt GTCTCTCAGC}$	AAGACAGAGT	AAGTTACTCT	TTTGTTAACA	700
ATCCTACCAG	TTATAATCAA	TTTAATTATG	GCCATGGGAG	ATGGAAAGAT	
TGGCAACAGC	GGGTACAAAA	AGATGTACGA	AATAAGCAAA	TAAGCTGTCA	800
TTCGTTAGAC	CTAGATTACT	TAAAAATAAG	TTTCACTGAA	AAAGGAAAAC	
AAGAAAATAT	TCAAAAGTGG	GTAAATGGTA	TATCTTGGGG	AATAGTGTAC	900
TATGGAGGCT	CTGGGAGAAA	GAAAGGATCT	GTTCTGACTA	TTCGCCTCAG	
AATAGAAACT	CAGATGGAAC	CTCCGGTTGC	TATAGGACCA	AATAAGGGTT	1000
TGGCCGAACA	AGGACCTCCA	ATCCAAGAAC	AGAGGCCATC	TCCTAACCCC	
TCTGATTACA	ATACAACCTC	TGGATCAGTC	CCCACTGAGC	CTAACATCAC	1100
TATTAAAACA	GGGGCGAAAC	TTTTTAGCCT	CATCCAGGGA	GCTTTTCAAG	
CTCTTAACTC	CACGACTCCA	GAGGCTACCT	CTTCTTGTTG	GCTTTGCTTA	1200
GCTTCGGGCC	CACCTTACTA	TGAGGGAATG	GCTAGAGGAG	GGAAATTCAA	
TGTGACAAAG	GAACATAGAG	ACCAATGTAC	ATGGGGATCC	CAAAATAAGC	1300
TTACCCTTAC	TGAGGTTTCT	GGAAAAGGCA	CCTGCATAGG	GATGGTTCCC	
CCATCCCACC	AACACCTTTG	TAACCACACT	GAAGCCTTTA	ATCGAACCTC	1400
TGAGAGTCAA	TATCTGGTAC	CTGGTTATGA	CAGGTGGTGG	GCATGTAATA	
CTGGATTAAC	CCCTTGTGTT	TCCACCTTGG	TTTTCAACCA	AACTAAAGAC	1500
TTTTGCGTTA	TGGTCCAAAT	TGTCCCCCGG	GTGTACTACT	ATCCCGAAAA	
AGCAGTCCTT	GATGAATATG	ACTATAGATA	TAATCGGCCA	AAAAGAGAGC	1600
			GATTGGGAGT		
GTGGGAACAG	GAACGGCTGC	CCTAATCACA	GGACCGCAAC	AGCTGGAGAA	1700
AGGACTTAGT	AACCTACATC	GAATTGTAAC	GGAAGATCTC	CAAGCCCTAG	

AAAAATCTGT CAGTAACCTG GAGGAATCCC TAACCTCCTT ATCTGAAGTG 1800 GTTCTACAGA ACAGAAGGGG GTTAGATCTG TTATTTCTAA AAGAAGGAGG GTTATGTGTA GCCTTAAAAG AGGAATGCTG CTTCTATGTA GATCACTCAG 1900 GAGCCATCAG AGACTCCATG AGCAAGCTTA GAGAAAGGTT AGAGAGGCGT CGAAGGGAAA GAGAGGCTGA CCAGGGGTGG TTTGAAGGAT GGTTCAACAG 2000 GTCTCCTTGG ATGACCACCC TGCTTTCTGC TCTGACGGGG CCCCTAGTAG TCCTGCTCCT GTTACTTACA GTTGGGCCTT GCTTAATTAA TAGGTTTGTT 2100 GCCTTTGTTA GAGAACGAGT GAGTGCAGTC CAGATCATGG TACTTAGGCA ACAGTACCAA GGCCTTCTGA GCCAAGGAGA AACTGACCTC TAGCCTTCCC 2200 AGTTCTAAGA TTAGAACTAT TAACAAGACA AGAAGTGGGG AATGAAAGGA TGAAAATGCA ACCTAACCCT CCCAGAACCC AGGAAGTTAA TAAAAAGCTC 2300 TAAATGCCCC CGAATTCCAG ACCCTGCTGG CTGCCAGTAA ATAGGTAGAA GGTCACACTT CCTATTGTTC CAGGGCCTGC TATCCTGGCC TAAGTAAGAT 2400 AACAGGAAAT GAGTTGACTA ATCGCTTATC TGGATTCTGT AAAACCGACT GGCACCATAG AA 2462

SEQ ID NO. 2: Translation of PERV-A env (1 letter code)

MHPTLSRRHLPIRGGK PKRLKIPLSFASIAWF LTLSITPQVNGKRLVD 48 SPNSHKPLSLTWLLTD SGTGININSTQGEAPL GTWWPELYVCLRSVIP GLNDQATPPDVLRAYG FYVCPGPPNNEEYCGN PODFFCKOWSCITSND 144 GNWKWPVSQQDRVSYS FVNNPTSYNQFNYGHG RWKDWQQRVQKDVRNK 192. QISCHSLDLDYLKISF TEKGKQENIQKWVNGI SWGIVYYGGSGRKKGS 240 VLTIRLRIETQMEPPV AIGPNKGLAEQGPPIQ EQRPSPNPSDYNTTSG 288 SVPTEPNITIKTGAKL FSLIQGAFQALNSTTP EATSSCWLCLASGPPY 336 YEGMARGGKFNVTKEH RDQCTWGSQNKLTLTE VSGKGTCIGMVPPSHQ 384 HLCNHTEAFNRTSESQ YLVPGYDRWWACNTGL TPCVSTLVFNQTKDFC 432 VMVQIVPRVYYYPEKA VLDEYDYRYNRPKREP ISLTLAVMLGLGVAAG 480 VGTGTAALITGPQQLE KGLSNLHRIVTEDLQA LEKSVSNLEESLTSLS 528 EVVLQNRRGLDLLFLK EGGLCVALKEECCFYV DHSGAIRDSMSKLRER 576 LERRRREREADQGWFE GWFNRSPWMTTLLSAL TGPLVVLLLLLTVGPC 624 LINRFVAFVRERVSAV QIMVLRQQYQGLLSQG ETDL\* 660

SEQ ID NO. 3: PERV-B.seq

GCATGCCTGC AGCAGTTGGT CAGAACATCC CCTTATCATG TTCTGAGGCT ACCAGGAGTG GCTGACTCGG TGGTCAAACA TTGTGTGCCC TGCCAGCTGG 100

TTAATGCTAA	TCCTTCCAGA	ATACCTCCAG	GAAAGAGACT	AAGGGGAAGC	
CACCCAGGCG	CTCACTGGGA	AGTGGACTTC	ACTGAGGTAA	AGCCGGCTAA	200
ATACGGAAAC	AAATATCTAT	TGGTTTTTGT	AGACACCTTT	TCAGGATGGG	
TAGAGGCTTA	TCCTACTAAG	AAAGAGACTT	CAACCGTGGT	GGCTAAAAAA	300
ATACTGGAGG	AAATTTTTCC	GAGATTTGGA	ATACCTAAGG	TAATCGGGTC	
AGACAATGGT	CCAGCTTTTG	TTGCCCAGGT	AAGTCAGGGA	CTGGCCAAGA	400
TATTGGGGAT	TGATTGGAAA	CTGCATTGTG	CATACAGACC	CCAAAGCTCA	
GGACAGGTAG	AGAGGATGAA	TAGAACCATT	AAAGAGACCC	TTACCAAATT	500
GACCACAGAG	ACTGGCATTA	ATGATTGGAT	AGCTCTCCTG	CCCTTTGTGC	
TTTTTAGGGT	TAGGAACACC	CCTGGACAGT	TTGGGCTGAC	CCCCTATGAA	600
TTGCTCTACG	GGGGACCCCC	CCCGTTGGTA	GAAATTGCTT	CTGTACATAG	
TGCTGATGTG	CTGCTTTCCC	AGCCTCTGTT	CTCTAGGCTC	AAGGCGCTCG	700
AGTGGGTGAG	GCAACGAGCG	TGGAAGCAGC	TCCGGGAGGC	CTACTCAGGA	
GAAGGAGACT	TGCAAGTTCC	ACATCGCTTC	CAAGTGGGAG	ATTCAGTCTA	800
TGTTAGACGC	CACCGTGCAG	GAAACCTCGA	GACTCGGTGG	AAGGGCCCTT	
ATCTCGTACT	TTTGACCACA	CCAACGGCTG	TGAAAGTCGA	AGGAATCTCC	900
ACCTGGATCC	ATGCATCCCA	CGTTAAGCTG	GCGCCACCTC	CCGACTCGGG	
GTGGAGAGCC	GAAAAGACTG	AGAATCCCCT	TAAGCTTCGC	CTCCATCGCC	1000
TGGTTCCTTA	CTCTAACAAT	AACTCCCCAG	GCCAGTAGTA	AACGCCTTAT	
ÁGACAGCTCG	AACCCCCATA	GACCTTTATC	CCTTACCTGG	CTGATTATTG	1100
ACCCTGATAC	GGGTGTCACT	GTAAATAGCA	CTCGAGGTGT	TGCTCCTAGA	
GGCACCTGGT	GGCCTGAACT	GCATTTCTGC	CTCCGATTGA	TTAACCCCGC	1200
TGTTAAAAGC	ACACCTCCCA	ACCTAGTCCG	TAGTTATGGG	TTCTATTGCT	
GCCCAGGCAC	AGAGAAAGAG	AAATACTGTG	GGGGTTCTGG	GGAATCCTTC	1300
TGTAGGAGAT	GGAGCTGCGT	CACCTCCAAC	GATGGAGACT	GGAAATGGCC	
GATCTCTCTC	CAGGACCGGG	TAAAATTCTC	CTTTGTCAAT	TCCGGCCCGG	1400
GCAAGTACAA	AGTGATGAAA	CTATATAAAG	ATAAGAGCTG	CTCCCCATCA	
GACTTAGATT	ATCTAAAGAT	AAGTTTCACT	GAAAAAGGAA	AACAGGAAAA	1500
TATTCAAAAG	TGGATAAATG	GTATGAGCTG	GGGAATAGTT	TTTTTATAAAT	
ATGGCGGGGG	AGCAGGGTCC	ACTTTAACCA	TTCGCCTTAG	GATAGAGACG	1600
GGGACAGAAC	CCCCTGTGGC	AGTGGGACCC	GATAAAGTAC	TGGCTGAACA	
GGGCCCCCG	GCCCTGGAGC	CACCGCATAA	CTTGCCGGTG	CCCCAATTAA	1700
CCTCGCTGCG	GCCTGACATA	ACACAGCCGC	CTAGCAACGG	TACCACTGGA	
TTGATTCCTA	CCAACACGCC	TAGAAACTCC	CCAGGTGTTC	CTGTTAAGAC	1800
				GCCATCAACT	
CCACCGACCC	TGATGCCACT	TCTTCTTGTT	GGCTTTGTCT	ATCCTCAGGG	1900
CCTCCTTATT	ATGAGGGGAT	GGCTAAAGAA	GGAAAATTCA	ATGTGACCAA	

AGAGCATAGA	AATCAATGTA	CATGGGGGTC	CCGAAATAAG	CTTACCCTCA	2000
CTGAAGTTTC	CGGGAAGGGG	ACATGCATAG	GAAAAGCTCC	CCCATCCCAC	
CAACACCTTT	GCTATAGTAC	TGTGGTTTAT	GAGCAGGCCT	CAGAAAATCA	2100
GTATTTAGTA	CCTGGTTATA	ACAGGTGGTG	GGCATGCAAT	ACTGGGTTAA	
CCCCCTGTGT	TTCCACCTCA	GTCTTCAACC	AATCCAAAGA	TTTCTGTGTC	2200
ATGGTCCAAA	TCGTCCCCCG	AGTGTACTAC	CATCCTGAGG	AAGTGGTCCT	
TGATGAATAT	GACTATCGGT	ATAACCGACC	AAAAAGAGAA	CCCGTATCCC	2300
TTACCCTAGC	TGTAATGCTC	GGATTAGGGA	CGGCCGTTGG	CGTAGGAACA	
GGGACAGCTG	CCCTGATCAC	AGGACCACAG	CAGCTAGAGA	AAGGACTTGG	2400
TGAGCTACAT	GCGGCCATGA	CAGAAGATCT	CCGAGCCTTA	GAGGAGTCTG	
TTAGCAACCT	AGAAGAGTCC	CTGACTTCTT	TGTCTGAAGT	GGTTCTACAG	2500
AACCGGAGGG	GATTAGATCT	GCTGTTTCTA	AGAGAAGGTG	GGTTATGTGC	
AGCCTTAAAA	GAAGAATGTT	GCTTCTATGT	AGATCACTCA	GGAGCCATCA	2600
GAGACTCCAT	GAGCAAGCTT	AGAGAAAGGT	TAGAGAGGCG	TCGAAGGGAA	
AGAGAGGCTG	ACCAGGGGTG	GTTTGAAGGA	TGGTTCAACA	GGTCTCCTTG	2700
GATGACCACC	CTGCTTTCTG	CTCTGACGGG	ACCCCTAGTA	GTCCTGCTCC	
TGTTACTTAC	AGTTGGGCCT	TGCTTAATTA	ATAGGTTTGT	TGCCTTTGTT	2800
AGAGAACGAG	TGAGTGCAGT	CCAGATCATG	GTACTTAGGC	AACAGTACCA	
AGGCCTTCTG	AGCCAAGGAG	AAACTGACCT	CTAGCCTTCC	CAGTTCTAAG	2900
ATTAGAACTA	TTAACAAGAC	AAGAAGTGGG	GAATGAAAGG	ATGAAAATGC	
AACCTAACCC	TCCCAGAACC	CAGGAAGTTA	ATAAAAAGCT	CTAAATGCCC	3000
CCGAATTCCA	GACCCTGCTG	GCTGCCAGTA	AATAGGTAGA	AGGTCACACT	
TCCTATTGTT	CCAGGGCCTG	CTATCCTGGC	CTAAGTAAGA	TAACAGGAAA	3100
TGAGTTGACT	AATCGCTTAT	CTGGATTCTG	TAAAACCGAC	TGGCACCATA	
GAAGAATTGA	TTACACATTG	ACAGCCCTAG	TGACCTATCT	CAACTGCAAT	3200
CTGTCACTCT	GCCCAGGAGC	CCACGCAGAT	GCGGACCTCC	GGAGCTATTT	
TAAAATGATT	GGTCCACGGA	GCGCGGGCTC	TCGATATTTT	AAAATGATTG	3300
GTCCACGGAG	CGCGGGCTCT	TCGATATTTT	AAAATGATTG	GTTTGTGACG	
CACAGGCTTT	GTTGTGAACC	CCATAAAAGC	TGTCCCGATT	CCGCACTCGG	3400
GGCCGCAGTC	CTCTACCCCT	GCGTGGTGTA	CGACTGTGGG	CCCCAGCGCG	
CTTGGAATAA	AAATCCTCTT	GCTGTTTGCA	TC		3482

SEQ ID NO. 4: Translation of PERV-B env (1 letter code)
MHPTLSWRHLPTRGGE PKRLRIPLSFASIAWF LTLTITPQASSKRLID 48
SSNPHRPLSLTWLIID PDTGVTVNSTRGVAPR GTWWPELHFCLRLINP 96
AVKSTPPNLVRSYGFY CCPGTEKEKYCGGSGE SFCRRWSCVTSNDGDW 144
KWPISLQDRVKFSFVN SGPGKYKVMKLYKDKS CSPSDLDYLKISFTEK 192

GKQENIQKWINGMSWG IVFYKYGGGAGSTLTI RLRIETGTEPPVAVGP 240
DKVLAEQGPPALEPPH NLPVPQLTSLRPDITQ PPSNGTTGLIPTNTPR 288
NSPGVPVKTGQRLFSL IQGAFQAINSTDPDAT SSCWLCLSSGPPYYEG 336
MAKEGKFNVTKEHRNQ CTWGSRNKLTLTEVSG KGTCIGKAPPSHQHLC 384
YSTVVYEQASENQYLV PGYNRWACNTGLTPC VSTSVFNQSKDFCVMV 432
QIVPRVYYHPEEVVLD EYDYRYNRPKREPVSL TLAVMLGLGTAVGVGT 480
GTAALITGPQQLEKGL GELHAAMTEDLRALEE SVSNLEESLTSLSEVV 528
LQNRRGLDLLFLREGG LCAALKEECCFYVDHS GAIRDSMSKLRERLER 576
RRREREADQGWFEGWF NRSPWMTTLLSALTGP LVVLLLLLTVGPCLIN 624
RFVAFVRERVSAVQIM VLRQQYQGLLSQGETD L\*

SEQ ID NO:5

WO 98/53104

TGGAAAGATTGGCAACAGCG (SEQ ID NO:5)

SEO ID NO:6

AGTGATGTTAGGCTCAGTGG (SEQ ID NO:6)

SEQ ID NO:7

TTCTCCTTTGTCAA--TTCCGG 3' (SEQ ID NO:7)

SEQ ID NO:8

TACTTTATCGGGTCCCACTG 3' (SEQ ID NO:8)

SEQ ID NO:9

TACTCTTTTGTTAACAATCCTA 3' (SEQ ID NO:9)

SEQ ID NO:10

TATTCTGAGGCGCGAATAGT 3' (SEQ ID NO:10)

SEQ ID NO:11

ATCCGTCGGCATGCATAATACGACTCAC (SEQ ID NO:11)

SEO ID NO:12

CGATTCAGTGCTGCTACAAC (SEQ ID NO:12)

SEQ ID NO:13

CCCTTATAACCTCTTGAGCG (SEQ ID NO:13)

SEQ ID NO:14

GTAATGCATGCTTCTATGGTGCCAGTCG (SEQ ID NO:14)

SEQ ID NO:15

CTCTACGCATGCGTGTGTACGACTGTG (SEQ ID NO:15)

SEQ ID NO:16

GTAATCGGGTCAGACAATGG (SEQ ID NO:16)

#### CLAIMS

- 1. An isolated nucleic acid probe, said probe being capable of hybridising to the PERV-B env gene under conditions in which said probe is substantially unable to hybridise to the PERV-A env gene.
- 2. An isolated nucleic acid probe according to claim 1 which is capable of hybridising to SEQ ID NO:3 or the complement thereof under conditions in which it is not capable of hybridising to SEQ ID NO:1 or the complement thereof.
- 3. An isolated nucleic acid probe according to claim 1 or 2 which is derived from the region of PERV-B derived from nucleotides 1000 to 2500 of the SEO ID NO. 3 isolate.
- 4. An isolated nucleic acid probe, said probe being capable of hybridising to the PERV-A *env* gene under conditions in which said probe is substantially unable to hybridise to the PERV-B *env* gene.
- 5. An isolated nucleic acid probe according to claim 4 which is capable of hybridising to SEQ ID NO:1 or the complement thereof under conditions in which it is not capable of hybridising to SEQ ID NO:3 or the complement thereof.
- 6. An isolated nucleic acid according to claim 4 or 5 which is derived from the region of PERV-A derived from nucleotides 300 to 1809 of the SEO ID NO:1 isolate.
- 7. An isolated nucleic acid probe according to any one of the preceding claims which is from 10 to 40 nucleotides in length.
- 8. A pair of primers suitable for conducting a polymerase chain reaction, at least one of said primers being a nucleic acid as defined in any one of claims 1 to 3.

- 9. A pair of primers suitable for conducting a polymerase chain reaction, at least one of said primers being a nucleic acid as defined in any one of claims 4 to 6.
- 10. A method of determining the subtype of a porcine endogenous retrovirus in a sample which contains or is suspected to contain one or both of the PERV-A and PERV-B subtypes, said method comprising probing said tissue with a nucleic acid probe as defined in any one of claims 1 to 7, or by conducting a polymerase chain reaction with a pair of primers as defined in claim 8 or 9, and determining whether or not said probe or pair of primers detects either of said subtypes.
- 11. A method according to claim 10 wherein retroviral material from said cells is amplified prior to probing or conducting said PCR.
- 12. A method according to claim 10 wherein the sample is cloned nucleic acid obtained from pig or human cells.
- 13. A method according to claim 10 or 11 wherein the sample comprises tissue which is primary porcine tissue.
- 14. A method according to claim 10 or 11 wherein the sample of is a human cell line which has been cultivated in the presence of a porcine cell line.
- 15. An antibody capable of binding to an epitope on the PERV-B env protein under conditions where said antibody is substantially unable to bind to the PERV-A env protein.
- 16. An antibody capable of binding to an epitope on the PERV-A env protein under conditions where said antibody is substantially unable to bind to the PERV-B env protein.
- 17. A method of detecting the presence of a pig endogenous

retrovirus in porcine or human tissue or cell lines which comprises bringing a sample of said tissue or cell line into contact with an antibody according to claim 15 or 16 and detecting whether or not said antibody binds to a retrovirus in the sample.

18. Use of a probe according to any one of claims 1 to 7 in a method of determining the subtype of a porcine endogenous retrovirus.

PERV-A	10 TCGAGTGGGTGAG				50 TACTCAGGA	60 GGAGACT
					AGA	
		-0.0		7.40		7.60
PERV-B	700	720		740	 	760
	70	80	90	100	110	120
PERV-A	TGCAĄGTTCCACA'					
		700		800		820
PERV-B		780 	G			
	130	140	150	160	170	180
PERV-A	GAAACCTCGAGAC'	TCGGTGGAAG	GACCTTATC?	rcgtacttttc	GACCACACCA	ACGGCTG
		840		860		880
PERV-B			c			
	190			220		
PERV-A	TGAAAGTCGAAGG	AATCCCCACC	rGGATCCATG(	CATCCCACGTT	TAAGCCGGCG	CCACCTC
		900		920		940
PERV-B		т			Т	
•	250			280		
PERV-A	CCGATTCGGGGTG	GAAAGCCGAAA	AAGACTGAAAA	ATCCCCTTAAC	GCTTCGCCTC(	CATCGCG
		960		980		1000
PERV-B	C	G				c
	310		330		350	
PERV-A	TGGTTCCTTACTC	TGTCAATAAC'	rcctcaagtt	AATGGTAAACC	CCTTGTGGA	CAGCCCG
		1020		1040		1060
PERV-B		.AA	cg.cc	.G.A	A.A	T
DEBM A	370 AACTCCCATAAAC	380	390 ~accrccrra	400 CTTACTGACT	410 CCGGTACAGG	420 TATTAAT
PERV-A	ANCICCCAIAAAC	CCLINICICI	LICCIGGI IA			
DODII D	CG	1080	m c c	1100	ጥ አ ር	1120 G C C
PERV-B	C G		1	A		

Fig. 1a

WO 98/53104 PCT/GB98/01428

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2/6 440 450 460 470 ATTAACAGCACTCAAGGGGAGGCTCCCTTGGGGACCTGGTGGCCTGAATTATATGTCTGC 1140 1160 G.A..T.....G...T.TT.....TAGA..C.........C.GC..T....510 520  ${\tt CTTCGATCAGTAATCCCTGGTCTCAATGACCAGGCCACACCCCCGATGTACTCCGTGCT}$ PERV-A ...C....TGA.T.A...C.C.G.T..------T...A.CC..G....AG. PERV-B 570 580 TACGGGTTTTACGTTTGCCCAGGACCCCCAAATAATGAAGAATATTGTGGAAATCCTCAG PERV-A 1260 1240  $\dots \texttt{T} \dots \texttt{C} \dots \texttt{TTGC} \dots \dots --- . \texttt{A} \dots \texttt{G} . \texttt{G} \dots \texttt{A} \dots \texttt{GA} \dots \dots \texttt{C} \dots \dots \texttt{GGG} . \texttt{T} \dots \texttt{GG}.$ PERV-B 620 630 640 650 660 610  ${\tt GATTTCTTTTGCAAGCAATGGAGCTGCATAACTTCTAATGATGGGAATTGGAAATGGCCA}$ 1320 PERV-B 680 690  ${\tt GTCTCTCAGCAAGACAGAGTAAGTTACTCTTTTGTTAACAATCCTACCAGTTATAATCAA}$ PERV-A 1400 1360 1380 A.....TC..G...C.G...AA.T...C.....C...-T...GG..C.GGCA.G.... 770 750 760 TTTAATTATGGCCATGGGAGATGGAAAGATTGGCAACAGCGGGTACAAAAAGATGTACGA PERV-A 1420 1440 AG.G..G.-AA.T..TAA....-A.G..C.GCT.CC..T.-A.ACTT.G.TT..C..-A.PERV-B 820 830 790 800 810 AATAAGCAAATAAGCTGTCATTCGTTAGACCTAGATTACTTAAAAATAAGTTTCACTGAA PERV-A

Fig. 1b

# SUBSTITUTE SHEET (rule 26)

G....-TT.C.-...AA.AAG.AA.ACAGG.A.A..T.C......A.GGT.-...C

GTGG

WO 98/53104 PCT/GB98/01428

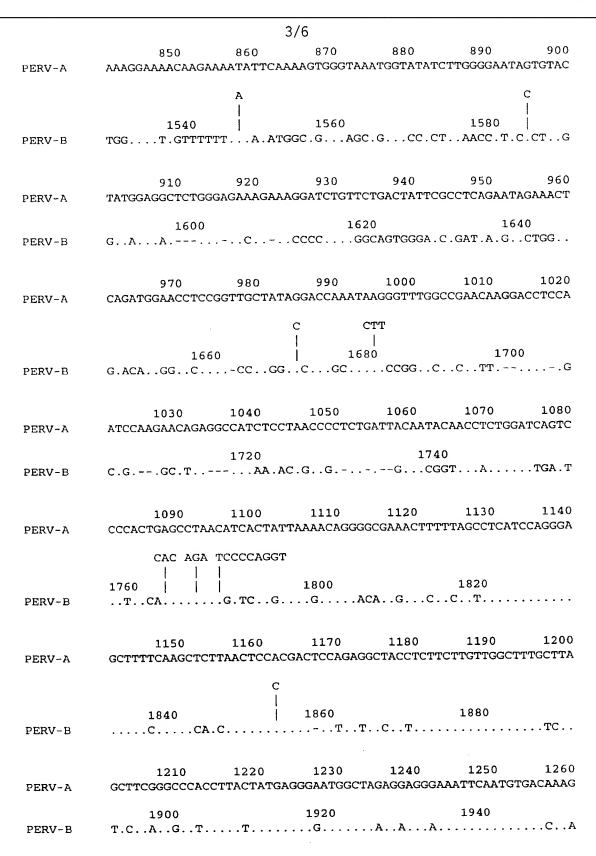


Fig. 1c

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25011				1300		
PERV-A	GAACATAGAGACC	AATGTACATG	GGGATCCCAA	AATAAGCITAC	CCTTACTGA	3GTTTCT
	1960		1980		2000	
PERV-B	GA.T.		GG.		C	AC
	1330	1340	1350	1360	1370	1380
PERV-A	GGAAAAGGCACCTC	GCATAGGGAT	GGTTCCCCCA'	TCCCACCAACA	CCTTTGTAA	CCACACT
	2020		2040		2060	
PERV-B	GGGA.	A.A	A.C		CT.	TAGT
				1420		
PERV-A	GAAGCCTTTAATC	GAACCTCTGA	GAGTCAATAT	CTGGTACCTGC	GTTATGACAG	GTGGTGG
	2080		2100		2120	
PERV-B	.TG.TT.A.G.G.	AGGA	A.AG	т.А	A	• • • • • •
PERV-A	1450 GCATGTAATACTG			1480		
PERV-A	GCATGTAATACTG	JATTAACCCC	.TIGIGITICC.	ACCITOGITI		
	2 <b>14</b> 0		2160		2180	С Т
PERV-B		.G	C	CA		C1
	1510	1520	1520	1540	1550	1560
PERV-A	TTTTGCGTTATGG					
	2200		2220		2240	
PERV-B	2200 CTC		2220 A	c		G
	1570	1580	1590	1600	1610	1620
PERV-A	GATGAATATGACT					
	2260		2280		2300	
PERV-B		c.g	CA	AG	т	c
	1630	1640	1650	1660	1670	1680
PERV-A	GTAATGCTCGGAT	TGGGAGTGG	TGCAGGCGTG	GGAACAGGAA	CGGCTGCCCT	AATCACA
	2320		2340		2360	
PERV-B		.AGAC	.C.TTA		.A	G
	•					
	1690	1700	1710			1740
PERV-A	GGACCGCAACAGC	TGGAGAAAG(	JACTTAGTAAC	CTACATCGAA	TTGTAACGGA	AGATCTC
	2380		2400		2420	
PERV-B	AG	.A	GG.G	GGCGG	CCA.GA	

Fig. 1d

Fig. 1e

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PERV-A	2230	2240	2250	2260	2270	2280
	TAACAAGACAAGA	AGTGGGGAA	FGAAAGGATGA	AAATGCAAC	CTAACCCTCCC	AGAACCC
PERV-B	2920	• • • • • • • • • • • • • • • • • • • •	2940		2960	
PERV-A	2290	2300	2310	2320	2330	2340
	AGGAAGTTAATAA	AAAGCTCTAA	ATGCCCCCGA	ATTCCAGACO	CCTGCTGGCTG	CCAGTAA
PERV-B	2980		3000	• • • • • • • • • • • • • • • • • • • •	3020	
PERV-A	2350	2360	2370	2380	2390	2400
	ATAGGTAGAAGGTO	CACACTTCCT	CATTGTTCCAG	GGCCTGCTAI	CCTGGCCTAA	GTAAGAT
PERV-B	3040		3060	· · · · · · · · · · · · · · · ·	3080	• • • • • • •
PERV-A	2410	2420	2430	2440	2450	2460
	AACAGGAAATGAGT	TGACTAATO	GCTTATCTGG.	ATTCTGTAAA	ACCGACTGGC	ACCATAG
PERV-B	3100		3120		3140	• • • • • •

Fig. 1f

# **PCT**





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C12Q 1/70, C07K 16/10, G01N 33/569 // C07K 14/15, C12N 15/48	A3	(43) International Publication Date: 26 November 1998 (26.11.98
(21) International Application Number: PCT/GB9 (22) International Filing Date: 18 May 1998 (18) (30) Priority Data: 9710154.7 16 May 1997 (16.05.97)  (71) Applicant (for all designated States except US): MERESEARCH COUNCIL [GB/GB]; 20 Park Cresces	8.05.9 C	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GI GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SI TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIP patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasia patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), Europea patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GI IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CI
don W1N 4AL (GB).  (72) Inventors; and (75) Inventors/Applicants (for US only): STOYE, Jonatha [GB/GB]; 28A Seaman Close, Park Street, St. Herts AL2 2NX (GB). WEISS, Robin, Anthony [C 25 Cyprus Avenue, Finchley, London N3 1SS (GB)	Albar GB/GF	s,
(74) Agents: BRASNETT, Adrian, H. et al.; Mewburn Ell House, 23 Kingsway, London WC2B 6HP (GB).	lis, Yo	rk

(54) Title: DETECTION OF RETROVIRAL SUBTYPES BASED UPON ENVELOPE SPECIFIC SEQUENCES

#### (57) Abstract

The present invention is based upon the finding that porcine endogenous retroviruses exist in two different subtypes, which we have termed PERV-A and PERV-B. The differences are reflected in sequence divergence in the envelope genes, and these differences may be used to provide nucleic acid and antibody probes which can distinguish between the two subtypes. This allows patterns of subtype transmission between cells, particularly porcine to human cells, to be monitored.

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## B. FIELDS SEARCHED

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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

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	vol. 389, October 1998, pages 681-82, XP002084123 see the whole document	
Р,А	WO 97 40167 A (Q ONE BIOTECH LTD ;IMUTRAN LTD (GB); GALBRAITH DANIEL NORMAN (GB);) 30 October 1997 see the whole document	1-18
Р,А	WO 97 21836 A (GEN HOSPITAL CORP) 19 June 1997 . see the whole document	1-18

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Date of the actual completion of theinternational search  12 November 1998	Date of mailing of the international search report  27/11/1998
Name and mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2  NL - 2280 HV Rijswijk  Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Osborne, H

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